

# Europäisches Patentamt European Patent Office Office européen des brevets



(11) EP 1 236 770 A1

(12)

### **EUROPEAN PATENT APPLICATION**

(43) Date of publication: 04.09.2002 Bulletin 2002/36

(51) Int Cl.7: C08L 23/10, C08L 23/16

(21) Application number: 01301873.4

(22) Date of filing: 01.03.2001

(84) Designated Contracting States:

AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU

MC NL PT SE TR

Designated Extension States:

AL LT LV MK RO SI

(71) Applicant: Borealis Technology Oy 06101 Porvoo (FI)

(72) Inventors:

Follestad, Arild
 3960 Stathelle (NO)

- Ommundsen, Espen 3970 Langesund (NO)
- Fosse, Knut
   3711 Skien (NO)
- (74) Representative: Campbell, Neil Boyd Frank B. Dehn & Co., 179 Queen Victoria Street London EC4V 4EL (GB)

### (54) Polyethylene compositions for rotomolding

(57) A polymer composition suitable for rotomoulding comprising

I) an ethylene homopolymer or copolymer with at least one other  $C_{3-10}$   $\alpha$ -olefin, having a melt flow rate of 0.5 to 30, a molecular weight distribution (Mw/Mn) of less than 4, an Mw of 50,000 to 110,000, a density of 0.940 g/cm<sup>3</sup> to 0.970 g/cm<sup>3</sup> and a melting point of 100 to 145°C;

OR

I) a propylene homopolymer or copolymer with at least one other  $C_{2-10}$   $\alpha$ -olefin, having a melt flow rate of 0.5 to 30, a molecular weight distribution (Mw/Mn) of less than 4, an Mw of 150,000 to 300,000, and a melting point of 100 to 170°C;

II) an ethylene homo or copolymer with at least one other  $C_{3-10}$   $\alpha$ -olefin, having a melt flow rate of within 40% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than

4. an Mw of within 30% of the Mw of component (I). a density of 0.880 g/cm<sup>3</sup> to 0.940 g/cm<sup>3</sup> said density being at least 0.010 g/cm<sup>3</sup> less than the density of component (I) and a melting point of at least 5°C less than that of component (I);

OR

II) a propylene homo or copolymer with at least one other  $C_{2-10}$   $\alpha$ -olefin having a melt flow rate of within 40% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4, an Mw of within 30% of the Mw of component (I), and a melting point of at least 10°C less than that of component (I).

### Description

10

15

20

30

35

50

[0001] This invention relates to the use of a particular polymer composition in for example, rotational moulding as well as to the polymer composition itself, to rotomoulding processes using the same and to rotomoulded articles made from the polymer composition. More specifically, the invention concerns the use of a polymer composition which comprises at least two components formed by single site catalysis having particular molecular weight distributions, comonomer compositions and densities.

[0002] Rotational moulding is a moulding process in which a particulate polymer, the moulding powder, is filled into a mould which is placed into an oven and rotated so that the polymer melts and coats the inside surface of the mould. In order to ensure that the moulded product is defect free, the moulding powder must have a relatively small particle size and should preferably be uniform in particle size and composition. Where, as is normal, the moulding powder has to contain colouring agents or other additives, e.g. stabilisers, the moulding powder is conventionally produced by grinding polymer pellets extruded from stabilised reactor grain powder to the correct particle size for rotation of moulding, usually this the colours or other additives being added in with the polymer pellets are mixed into the ground and moulding powder.

[0003] A wide variety of articles may be prepared by rotational moulding. In particular rotational moulding is used in the manufacture of large objects such as liquid containers, e.g. tanks, boats, as well as in a large number of household areas, e.g. in the manufacture of toys.

[0004] The nature of the polymer rotomoulded depends very much on the nature of the article to be made. For example, if a chemical tank is being made, then the polymer used should be one which is not degraded by the chemical and one which has particular mechanical properties so that the container does not break under stress. Polymers used for the manufacture of toys must be completely non-toxic and again must be strong to prevent breakage. Articles for outdoor use such as boats must also be resistant to degradation from by the elements, e.g. sunlight, rain, frost or seawater. The mechanical properties of the rotomoulding powders are therefore critical.

[0005] Another important property is rheology and it is also critical that this is favourable. Rheology is a measure of non-Newtonian solid flow and it is crucial that flow be within certain limits to ensure that product properties are ideal.

[0006] Moreover, when making objects where a well defined shape is required, it is also desired that the eventual rotomoulded product does not warp, i.e. that the sides of a product remain undistorted.

[0007] A variety of polymers may be successfully rotomoulded although homo and copolymers of ethylene and homo and copolymers of propylene may in particular be mentioned. However, the nature of the catalyst used to make the polymer has a significant bearing on the rotomouldability of the polymer.

[0008] In rotomoulding, polymers produced from single site catalysts give rise to products having excellent mechanical properties and enable rotomoulding to be carried out over a much shorter period of time. Polymers produced from single site catalysts tend to have a narrow molecular weight distribution and copolymers produced from single site catalysts tend to have narrow comonomer distribution. These properties gives rise to increased environmental stress cracking resistance and improvements in other mechanical properties.

**[0009]** However, the narrow distribution of comonomers as compared to a Ziegler-Natta produced polymer, results in a much narrower melting and crystallisation behaviour. The sharp melting behaviour makes the polymer very sensitive to processing temperature and, without wishing to be limited by theory, it is believed that this causes severe warpage in rotomoulding products. Hence, the mechanical property benefits of using a polymer made by single site catalysis are offset by increased warpage.

[0010] Polymers produced using Ziegler-Natta catalysts have much broader melting/crystallisation windows than polymers made by single site catalysis and hence tend to produce rotomoulding products with much less warpage. However, due to the broad comonomer distribution and broad molecular weight distribution mechanical properties, especially ESCR and rheology are not so favourable.

[0011] There still remains therefore, the need to find a polymer suitable for rotational moulding that can give rise to products having both low warpage and excellent rheological and mechanical properties.

[0012] It has now been surprisingly found that by forming, e.g. blending, a particular mix of polymers, preferably made by single site catalysis a polymer composition may be produced which not only has excellent mechanical and rheological properties but also does not warp after rotomoulding since its processing window is broadened. A blend of polymers having, depending on the monomers involved, similar molecular weights and similar melt flow rates but different densities, melting points or comonomer distributions has surprisingly been found to give rise to a composition which shows an overall narrow molecular weight distribution and hence excellent mechanical and rheological properties and has a broadened processing window which eliminates warpage normally associated with single site materials.

[0013] Hence, viewed from one aspect the invention provides a polymer composition suitable for rotomoulding comprising

I) a first ethylene homo or copolymer with at least one other  $C_{3-10}$   $\alpha$ -olefin, having a melt flow rate (MFR) of 0.5 to

30, preferably 3 to 15, especially 6 to 8, a molecular weight distribution (Mw/Mn) of less than 4, preferably less than 3.5, especially less than 3, an Mw of 50,000 to 110,000, a density of 0.940 g/cm<sup>3</sup> to 0.970 g/cm<sup>3</sup> and a melting point of 100 to 145°C;

OR

5

10

30

35

45

- I) a propylene homo or copolymer with at least one other  $C_{2-10}$   $\alpha$ -olefin, having a melt flow rate of 0.5 to 30. preferably 3 to 15, especially 6 to 8, a molecular weight distribution (Mw/Mn) of less than 4, preferably less than 3.5, especially less than 3, an Mw of 150,000 to 300,000, and a melting point of 100 to 170°C: and
- II) a second ethylene homo or copolymer with at least one other C<sub>3-10</sub> α-olefin, having a melt flow rate of within 40%, preferably 20% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4, an Mw of within 30%, preferably 20% of the Mw of component (I), a density of 0.880 g/cm³ to 0.940 g/cm³ said density being at least 0.010 g/cm³ less than the density of component (I) and a melting point of at least 5°C, preferably at least 10°C less than that of component (I):
- II) a propylene homo or copolymer with at least one other  $C_{2-10}$   $\alpha$ -olefin having a melt flow rate of within 40%, preferably 20% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4. an Mw of within 30%, preferably 20% of the Mw of component (I), and a melting point of at least 10°C less than that of component (I).
- [0014] Viewed from another aspect the invention provides a polymer composition suitable for rotomoulding comprising as hereinbefore described wherein said composition has a molecular weight distribution (Mw/Mn) of less than 4, an Mw of within 30%, preferably 20% of the Mw of component (I), and an Mn within 30%, preferably 20% of the Mw of component (I).
  - [0015] In an especially preferred embodiment, the invention provides a polymer composition suitable for rotomoulding comprising
    - I) an ethylene homopolymer having a melt flow rate of 0.5 to 30, preferably 3 to 15, especially 6 to 8, a molecular weight distribution (Mw/Mn) of less than 4, preferably less than 3.5, especially less than 3, an Mw of 50,000 to 110,000, a density of 0.940 g/cm<sup>3</sup> to 0.970 g/cm<sup>3</sup> and a melting point of 100 to 145°C; and
    - II) an ethylene copolymer with at least one  $C_{3\cdot10}$   $\alpha$ -olefin having a melt flow rate of within 40%, preferably 20% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4, an Mw of within 30%, preferably 20% of the Mw of component (I), a density of 0.880 g/cm<sup>3</sup> to 0.940 g/cm<sup>3</sup>, preferably 910 g/cm<sup>3</sup> to 0.930 g/cm<sup>3</sup> said density being at least 0.010 g/cm<sup>3</sup> less than the density of component (I) and a melting point of at least 10°C less than that of component (I).
    - [0016] Viewed from another aspect the invention provides the use of a polymer composition as hereinbefore described in rotomoulding.
- [0017] Viewed from yet another aspect the invention provides a process for the preparation of an article comprising rotomoulding a composition as herein before defined.
  - [0018] Viewed from yet another aspect the invention provides an article comprising a polymer composition as here-inbefore described, especially a rotomoulded article.
  - [0019] Viewed from yet another aspect the invention provides a process for the preparation of a polymer composition as hereinbefore described wherein said composition is produced in at least two reactors in cascade or parallel, e.g. two slurry phase reactors or two gas phase reactors, especially a slurry phase followed by gas phase reactor.
  - [0020] Viewed from another aspect the invention provides a a process for the preparation of a polymer composition as hereinbefore described comprising blending components (I) and (II) in conventional blending apparatus, preferably a micropellet extruder.
- [0021] Viewed from another aspect the invention provides a a process for the preparation of a polymer composition as hereinbefore described said process comprising the use of at least two catalysts, preferably metallocene catalysts, e.g. a dualsite catalyst preferably a dualsite metallocene catalyst.
  - [0022] Unless otherwise stated densities are measured according to ISO 1183-1987(E). MFR is measured according to ISO 1133-1997 (D -for PE/M for PP). The melting point of polyethylene is measured by heating the polymer from room temperature to 200°C at a heat rate of 10°C/min. Thee polymer is maintained at 200°C for 5 mins and then cooled to -10°C at a cool rate of 10°C/min and maintained at -10°C for 1 minute. The polymer is then heated to 200°C at a heat rate of 10°C/min and the melting point is taken on this second heat run. For polypropylene the procedure is identical except that heating takes place to 225°C and cooling is effected to 20°C. GPC analyses were carried out

under the following conditions:

5

10

15

25

30

35

40

45

50

Equipment: Waters 150 CV plus no. 1115

Detector: Refractive Index (RI) and Viscosity detector Calibration: Narrow molecular weight distribution PS 1. Columns: 3 x HT6E styragel from Waters (140°C).

[0023] Components (I) and (II) may be copolymers of ethylene or propylene with at least one other  $C_{2-10}$   $\alpha$ -olefin. Suitable comonomers include ethylene, propylene, 1-butene, 1-hexene, 1-octene etc. Clearly, ethylene is only a suitable comonomer when the major monomer is propylene and propylene is only a suitable comonomer where the major monomer is ethylene. Diolefins may also be employed as comonomers especially those having two terminal double bonds, e.g. butadiene. In a preferred embodiment, where component (I) or (II) is a copolymer, it is a copolymer of ethylene with octene, butene or hexene, especially butene or hexene.

[0024] Component (I) or (II) may also be a homopolymer of propylene having no or only a few crystallinity disrupting units, e.g. less than 5 units per 100 propylene linkages. By crystallinity disrupting units it is meant a unit that disrupts the regular structure of the polymer, i.e. an atatic unit in syndiotactic or isotactic propylene polymers. In one embodiment, both components (I) and (II) should be propylene homopolymers in which component (I) is preferably a syndiotactic or isotactic propylene component and component (II) is preferably an amorphous propylene homopolymer. In a less preferred embodiment component (I) may be a polypropylene homopolymer and component (II) may be a propylene copolymer.

[0025] In another less preferred embodiment both components may be ethylene homopolymers in which component (II) comprises a high degree of short chain branching.

[0026] In a final and most preferred embodiment, component (I) may be an ethylene homopolymer and component (II) may be an ethylene copolymer.

[0027] The polymer components (I) and (II) should preferably have similar melt flow rates, i.e. the melt flow rate of component (II) should not differ from the melt flow rate of component (I) by greater than 40%, e.g. 20%, preferably no more than 10%, especially no more than 5%. The melt flow rate of the components should be in the range 0.5 to 30, preferably 1 to 20, more preferably 3 to 15, e.g. 4 to 10, especially 6 to 8, most especially about 6. In a highly preferred embodiment both components (I) and (II) have a MFR of about 6.

[0028] The melt flow rate of the entire composition should also be in the range 0.5 to 30, preferably 1 to 20, more preferably 2 to 15, e.g. 4 to 10, especially 6 to 8, most especially about 6.

[0029] The molecular weight distribution (MWD) of both components should be approximately the same, e.g. within 10%, and the MWD must be narrow, e.g. an (Mw/Mn) of less than 4, preferably less than 3.5, especially less than 3. The MWD of the entire composition should also preferably be less than 4, especially less than 3.5.

[0030] Whilst the Mw and Mn ranges may vary within wide limits the Mw/Mn ratio remains low, i.e. less than 4. In a preferred embodiment the Mw and Mn of both components are also similar. For a polyethylene homopolymer or copolymer suitable Mw values are in the range 50000 to 110000, especially 65000 to 85000. For a propylene homopolymer or copolymer suitable Mw values are Mw of 150,000 to 300,000. The Mw of component (II) should be within 30%, preferably 20% of the Mw of component (I).

[0031] For compositions based on ethylene, the densities of the two components should be different, i.e. component (II) should have a density at least 0.010 g/cm<sup>3</sup>, especially 0.020 g/cm<sup>3</sup> different from component (I). For ethylene homo and copolymers, preferably component (I) should have a density in the range 0.940 to 0.970 g/cm<sup>3</sup> and component (II) should have a density in the range 0.910 g/cm<sup>3</sup> to 0.930 g/cm<sup>3</sup>, said density preferably being at least 0.010 g/cm<sup>3</sup> less than the density of component (I).

[0032] For compositions based on ethylene, the density of the entire composition is preferably in the range 0.925 to 0.50 g/cm<sup>3</sup>, preferably 930 to 0.940 g/cm<sup>3</sup>.

[0033] The two components must also have different crystallinity properties, i.e. components (I) and (II) should have different melting points. This may be achieved by providing copolymers with differing comonomer contents or for a composition containing only homopolymers by providing a polymer having differing numbers of crystallinity disrupting units (e.g. short chain branches).

[0034] Where component (I) is an ethylene homo or copolymer it may have a melting point of 100 to 145°C. Where component (I) is a propylene copolymer or propylene homopolymer it should have a melting point of 100 to 170°C.

[0035] Component (II) should have a melting point which differs from that of component (II) by at least 5°C, preferably at least 10°C, especially at least 20°C. In the case of a propylene homopolymer, component (II) may also be amorphous and may therefore have no defined melting point.

[0036] Thus an especially preferred composition according to the invention is a composition in which component (I) is an ethylene homopolymer having an MFR of 6 to 8, a molecular weight distribution of less than 3, an Mw of 65,000 to 100,000, an Mn of 20,000 to 60,000, a density of 0.945 to 0.970 g/cm<sup>3</sup> and a melting point of 125 to 135°C; and component (II) is an ethylene copolymer with hexene having an MFR of 6 to 8, a molecular weight distribution of less

than 3, an Mw of 65,000 to 100,000, an Mn of 20,000 to 60.000. a density of 0.910 to 0.940 g/cm<sup>3</sup> and a melting point of 100 to 125°C.

[0037] In order to prepare the required polymer for rotomoulding, components (I) and (II) may be blended using conventional blending or compounding technology. The components (I) and (II) may be mixed in any convenient ratio to ensure that the desired properties are obtained. Preferably however, the ratio of component (I) to (II) is from 95:5 to 5:95, preferably 9:1 to 1:9, especially 4:1 to 1:4, more especially 1:2 to 2:1.

[0038] Components (I) and (II) may also be used in conjunction with other polymers in the blend such as rotomoulding polymer grades and some Ziegler-Natta polymers. Moreover, it is within the scope of the invention to use a further polymer component (III) which also has a MWD and MFR similar to components (I) and (II) but has a still different component distribution hence producing a multimodal component distribution.

[0039] The polymer composition described above gives rise to rotomoulded articles with excellent mechanical and rheological properties and low warpage.

[0040] The components are preferably produced using a single site catalyst, e.g. metallocene catalyst or potentially a dualsite catalyst. However, where component (I) or (II) is a homopolymer Ziegler-Natta catalysis may be employed. This is not however, preferred. Suitable metallocene catalysts for use in the invention may be any conventional metallocene catalyst. As used herein, the term metallocene is used to refer to all catalytically active metal:η-ligand complexes in which a metal is complexed by one, two or more open chain or closed ring η-ligands. The use of bridged bis- $\eta$ -ligand metallocenes, single  $\eta$ -ligand "half metallocenes", and bridged  $\eta$ - $\sigma$  ligand "scorpionate" metallocenes is particularly preferred. The metal in such complexes is preferably a group 4A, 5A, 6A, 7A or 8A metal or a lanthanide or actinide, especially a group 4A, 5A or 6A metal, particularly Zr, Hf or Ti. The η-ligand preferably comprises an η<sup>4</sup> or η<sup>5</sup> open chain or an η<sup>5</sup>-cyclopentadienyl ring, optionally with a ring or chain carbon replaced by a heteroatom (e.g. N, B, S or P), optionally substituted by pendant or fused ring substituents and optionally linked by bridge (e.g. a 1 to 4 atom bridge such as (CH<sub>2</sub>)<sub>2</sub>, C(CH<sub>3</sub>)<sub>2</sub> or Si(CH<sub>3</sub>)<sub>2</sub>) to a further optionally substituted homo or heterocyclic cyclopentadienyl ring. The ring substituents may for example be halo atoms or alkyl groups optionally with carbons replaced by heteroatoms such as O, N and Si, especially Si and O and optionally substituted by mono or polycyclic groups such as phenyl or naphthyl groups. Suitable η-ligands, include those of formula II discussed above. Examples of such homo or heterocyclic cyclopentadienyl ligands are well known in the art (see e.g. EP-A-416815, WO96/04290, EP-A-485821, EP-A-485823, US-A-5276208 and US-A-5145819).

[0041] Besides the  $\eta$ -ligand, the metallocene complex used according to the invention may include other ligands; typically these may be halide, hydride, alkyl, aryl, alkoxy, aryloxy, amide, carbamide or other two electron donor groups. Any hydrocarbyl ligand here will generally contain up to 20 carbons, preferably up to 10 carbons, e.g. up to 6 carbons. [0042] Metallocene catalysts are conventionally employed in the presence of a cocatalyst. Suitable cocatalysts are well known and include alkyl metal compounds, in particular alumoxanes. Suitable alumoxanes include  $C_{1-10}$  alkyl alumoxanes, e.g. methyl alumoxane (MAO) and isobutyl alumoxanes (e.g. tetra and hexaisobutyl alumoxane, TIBAO and HIBAO), especially MAO. Alumoxane co-catalysts are described by Hoechst in WO-A-94/28034. These are considered cyclic or cage like oligomers having up to 40, preferably 3 to 20, -[Al(R")O]- repeat units (where R" is hydrogen,  $C_{1-10}$  alkyl, preferably methyl, or  $C_{6-18}$  aryl or mixtures thereof).

[0043] If desired the metallocene or metallocene/cocatalyst mixture may be used in unsupported form or it may be precipitated and used as such. However the metallocene or its reaction product with the cocatalyst is preferably introduced into the polymerization reactor in supported form, e.g. impregnated into a porous particulate support, as is well known in the art.

[0044] The particulate support material used is preferably an organic or inorganic material, e.g. a polymer (such as for example polyethylene, polypropylene, an ethylene-propylene copolymer, another polyolefin or polystyrene or a combination thereof). Such polymeric supports may be formed by precipitating a polymer or by a prepolymerization, eg of monomers used in the polymerization for which the catalyst is intended. However, the support is especially preferably a metal or pseudo metal oxide such as silica, alumina or zirconia or a mixed oxide such as silica-alumina, in particular silica, alumina or silica-alumina.

[0045] Especially preferably the support is a porous material so that the metallocene may be loaded into the pores of the support, e.g. using a process analogous to those described in WO94/14856 (Mobil), WO95/12622 (Borealis) and WO96/00243 (Exxon). The particle size is not critical but is preferably in the range 5 to 200  $\mu$ m, more preferably 10 to 80  $\mu$ m.

[0046] Before loading, the particulate support material is preferably calcined, i.e. heat treated, preferably under a non-reactive gas such as nitrogen. This treatment is preferably at a temperature in excess of 100°C, more preferably 200°C or higher, e.g. 200-800°C, particularly about 300°C. The calcination treatment is preferably effected for several hours, e.g. 2 to 30 hours, more preferably about 10 hours.

[0047] A cocatalyst, e.g. an alumoxane or an ionic catalyst activator (such as a boron or aluminium compound, especially a fluoroborate) may also be mixed with or loaded onto the catalyst support material. This may be done subsequently or more preferably simultaneously to loading of the metallocene, for example by including the cocatalyst

10

20

40

50

in the solution of the metallocene or, by contacting the metallocene loaded support material with a solution of the cocatalyst or catalyst activator, e.g. a solution in an organic solvent. Alternatively however, any such further material may be added to the metallocene loaded support material in the polymerization reactor or shortly before dosing of the catalyst material into the reactor.

[0048] In this regard, as an alternative to an alumoxane it may be preferred to use a fluoroborate catalyst activator, especially a  $B(C_6F_5)_3$  or more especially a  $B(C_6F_5)_4$  compound, such as  $C_6H_5N(CH_3)_2H:B(C_6F_5)_4$  or  $(C_6H_5)_3C:B(C_6F_5)_4$ . Other borates of general formula (cation+)<sub>a</sub> (borate-)<sub>b</sub> where a and b are positive numbers, may also be used. [0049] Both components (I) and (II) may be prepared simultaneously using a dualsite catalyst, i.e. a catalyst carrying two active metallocene sites on a single support, one site designed to give a component (I) and the other designed to give component (II).

[0050] The polymerisation is typically conducted in the presence of a diluent. As a diluent, a linear, branched or cyclic saturated hydrocarbon such as butane, propane, pentane, hexane, heptane, octane, cyclohexane or methylcyclohexane may be used.

[0051] Polymerisation to produce the polymer for use in the invention may take place in the slurry, solution or gas phase. Slurry phase polymerisation can be conducted under standard slurry conditions.

[0052] For slurry reactors, the reaction temperature will generally be in the range 60 to 110°C (e.g. 80-110°C), the reactor pressure will generally be in the range 5 to 80 bar (e.g. 25-65 bar), and the residence time will generally be in the range 0.3 to 5 hours (e.g. 0.5 to 2 hours). The diluent used will generally be an aliphatic hydrocarbon having a boiling point in the range -70 to +100°C, especially isobutane or propane.

[0053] For solution phase reactors, the reaction temperature used will generally be in the range 130 to 270°C, the reactor pressure will generally be in the range 20 to 400 bar and the residence time will generally be in the range 0.1 to 1 hour. The solvent used will commonly be a hydrocarbon with a boiling point in the range 80-200°C.

[0054] For gas phase reactors, the reaction temperature used will generally be in the range 60 to 115°C (e.g. 70 to 110°C), the reactor pressure will generally be in the range 10 to 25 bar, and the residence time will generally be 1 to 8 hours. The gas used will commonly be a non-reactive gas such as nitrogen together with monomer (e.g. ethylene). [0055] In order to ensure that copolymer particles are in the correct size for rotational moulding the products of any polymerisation reaction may be converted to powder form or pelletized to a particle size of approximately 0.1 to 0.5 mm, preferably 0.3 mm using standard technology. Hence, suitably sized pellets may be prepared by grinding.

**[0056]** Alternatively micropellets may be produced using the technique described in WO 00/35646 which is hereby incorporated by reference. By this method a mixture of polyolefin and optionally at least one additive is extruded in melt form through a die and pelletised to give particles having a particular size distribution. The particles are then dried to very low levels of moisture to improve rotomouldability.

[0057] Alternatively, the polymerisation, using a dualsite or multisite catalyst, can be set up such that the reactor powder is suitable for use without further manipulation. In one embodiment of the invention rotomoulding may be carried out by combining polymer powder with a masterbatch of UV-stabiliser-loaded polyolefin powder in line with the teaching of WO00/11065 which is hereby incorporated by reference.

[0058] The polymer powder or pellets can comprise any standard additives e.g. one or more selected from colouring agents, stabilisers, antioxidants, UV-absorbers, anti-static agents, lubricants and fillers.

[0059] Rotational moulding may take place under standard conditions. The polymer powder is placed in the mould which is then transferred to an oven and rotated, preferably about two axes to distribute the polymer powder over the hot surfaces of the mould. The heating cycle is continued until all of the powder has melted and formed a thick, continuous layer within the mould. The mould is then removed from the oven and cooled until the resin has solidified. The moulded part is then removed.

[0060] The length of time which the mould must be heated depends on the nature of the article being moulded, the amount of resin present and the temperature of the oven. Typical rotomoulding temperatures are 230°C to 350°C, more particularly 260°C to 320°C. Heating time is chosen such that the inner air temperature in the mould is 160°C to 300°C, more preferably 170°C to 250°C. This temperature can be measured using a Rotolog® or similar equipment to monitor the temperature or it may be chosen based on previous experience. Cooling may be carried out under a stream of air, water spray or mist or simply in ambient air at room temperature. A combination of these methods may also be employed. Preferably cooling is achieved using a combination of blown air followed by ambient air or just blown air. Cooling times are normally of similar magnitude to heating times or slightly longer. Slow cooling reduces the amount of warpage present in a rotomoulded article however, it is a purpose of the invention to provide polymer compositions which can be cooled more rapidly without increases in warpage compared to conventional single site polymers. The moulded tank may be removed from its mould at any convenient time although it is preferred if it is removed when it has cooled to a temperature of 60°C to 100°C.

[0061] The skilled artisan is able to manipulate the temperature, time and rotation speed/ratio within a rotomoulding apparatus to ensure that well-formed moulded articles are produced.

[0062] Particularly preferr d rotomoulding conditions are Rotation Speed 9/1.4 RPM; heating for 13 minutes in oven

20

35

at 270°C; fan assisted cooling for 10 minutes followed by ambient air cooling for 6 minutes.

[0063] The polymer composition of the invention may also have utility outside the field of rotomoulding. It is envisaged that the composition may give benefits in thermoforming due to the broadened processing window which the polymers exhibit. In particular polymer compositions of this type may be useful in the replacement of PVC. The polymer compositions may also be useful in film and injection moulding applications and this forms a yet further aspect of the invention.

[0064] The invention will now be further illustrated with reference to the following non-limiting examples and figures.

### Example 1

20

25

30

35

[0065] Two polymer components A and B were prepared as described in Annex 1 below. The catalyst used was made from (nBu-Cp)ZrCl2 and MAO impregnated on a support of calcined silica. Polymerisations took place in a bench scale semibatch reactor with hydrogen premixed in Component A is an ethylene homopolymer made from a single site catalyst having a density of 0.957 g/cm³, an Mw of 77000, an Mn of 28000 giving a Mw/Mn of 2.7, a melting point of 132°C and a MFR of 6.

[0066] Component B is an ethylene/hexene copolymer having a density of 0.923 g/cm³, a Mw of 67000 and Mn of 31000 giving an Mw/Mn of 2.1, an MFR of 8 and a melting point of 120°C. These components were compounded together, optionally with further polyethylene reactor powders, RP1 and RP2 having a density of 0.934 g/cm³ and an MFR of 6 or a density of 0.939 g/cm³ and an MFR of 6 respectively. The polymer blend was completed by the addition of an antioxidant and a slip agent.

[0067] The polymer blends prepared are shown in Table 1 below:

To	h	$\sim$	-

Sample	Comp. A	RP1	RP2	Comp. B
934Y	18%	25%	57%	-
934A	64%	-	-	36%
934B	32%	50%	-	18%
934C	16%	75%	-	9%
934D	8%	87.5%	-	4.5%
934Ref	-	100%	-	-

[0068] All blends were made to a total of 4 kg which resulted in approximately 3.8 kg of granules. These blends were then ground leaving at least 2.8 kg of powder. The overall density of each blend was 0.934 g/cm<sup>3</sup>, except 934Y which had a density of 0.940 g/cm<sup>3</sup>.

[0069] In addition, some commercially available polymer grades were readied for rotomoulding.

[0070] RM 8403, an ethylene/hexene copolymer available from Borealis A/S, is a polymer produced from a metal-locene catalyst having the following properties: Mw=75000, Mn=34000, MWD=2.2, MFR 6, density 940 g/cm<sup>3</sup>, melting point 125°C, heat of fusion 194 J/g, cryst Temp 110°C, heat of cryst -158 J/g.

[0071] RM 8343, an ethylene/hexene copolymer available from Borealis A/S, is a polymer produced from a metal-locene catalyst having the following properties: Mw 76000, Mn=34000, MWD=2.2 MFR 6, density 934 g/cm³, melting point 123°C, heat of fusion 176 J/g, cryst Temp 108°C, heat of cryst -156 J/g.

[0072] ME8152, an ethylene/butene copolymer available from Borealis A/S, is a polymer produced from a Ziegler-Natta catalyst having the following properties: Mw 101000, Mn 25000, MWD 4.1, MFR 3.5, density 934 g/cm<sup>3</sup>, melting point 125°C, heat of fusion 180 J/g, cryst Temp 110°C.

[0073] The properties of each blend are further explained in the table below

	934ref	934A	934B	934C	934D	934Y
Density	935.1	934.9	935.3	935.3	934.6	940.6
Melt pt	123.4	127	125	124.4	123.7	127
heat of fusion	163	163	164	164	160	182
cryst T	109	112.3	110.6	110	109.6	112.6
heat of cryst	-155	-151	-153	-154	-148	-173

### Example 2

10

25

30

35

45

50

55

[0074] Samples were rotomoulded under one or more of the following sets of conditions to form boxes.

### R tom ulding S rial 1

[0075] No preheating: Rotation Speed 9/1.4 RPM; heating for 13 minutes in oven at 270°C; fan assisted cooling for 10 minutes followed by 6 minutes ambient cooling in the absence of a fan. 700 g of polymer employed; Max mould temperature 227°C, mould temperature at start 35°C.

### **Rotomoulding Serial 3**

[0076] Mould was preheated to 60°C; Rotation Speed 4/2; heating for 13 minutes in oven at 270°; fan assisted cooling for 30 minutes. 700 g of polymer employed; Max mould temperature 227°C, mould temperature at start 60°C. [0077] The boxes resulting from the rotomoulding were cubes having sides of approximately 20 cm. The edges of the boxes were trimmed.

### Example 3

[0078] Warpage of the rotomoulded boxes was measured on five of the six cube walls (not the top wall). A ruler with a micrometer was used diagonally on each side of the box. Results are depicted in Table 2 relative to results achieved with ME8152.

Table 2

Вох	Average Warpage
ME8152 - Roto Serial 3	100%
ME8152 - Roto Serial 1	66%
RM8343 - Roto Serial 3	143%
RM8343 - Roto Serial 1	92%
934A - Roto Serial 3	72%
934A - Roto Serial 1	38%

[0079] These results are graphically depicted in Figure 1.

### Example 4

[0080] Thickness variation in the walls of the boxes was measured by taking five wall thickness measurements from each side of the boxes. Average thickness variation is standard deviation for all points of each box relative to ME8152. The results are depicted in Table 3 below.

Table 3

Box	Thickness variation
ME8152 - Roto Serial 3	100%
ME8152 - Roto Serial 1	38%
RM8343 - Roto Serial 3	115%
RM8343 - Roto Serial 1	33%
934A - Roto Serial 3	53%
934A - Roto Serial 1	59%

[0081] The data collected to obtain these results is shown in the attached Annex 2 in which 9\_1.4 std represents rotomoulding Serial 1 and 4\_2 full cool represents rotomoulding serial 3.

### Discussion

[0082] Blends with the broadest comonomer distribution have significantly less warpage and thickness variation than the reference materials. Moreover, an analysis of the crystal lattice structure of 934A reveals a much finer crystal structure which should make the material more robust to morphological changes. The appearance of the boxes made with 934A was also improved and fewer air bubbles are formed in the rotomoulded article.

### Example 5

10 [0083] Mechanical Properties. The following tests were employed to test the mechanical properties of the boxes produced.

Tensile Modulus: ISO527-1 (1993)

Instrument Falling Weight (IFW) ISO 6603-2: 1989

[0084] Circular disks with diameter 60 mm are used with a hemispherical striker of mass 10 kg and 20 mm diameter. Falling height 1 m at a velocity of 4.4m/s at -20°C. Rupture was ductile.

ESCR - ASTM D1693-97/ISO 1872-2: 1997

[0085] Standardised specimens are notched and stressed before being lowered into a solution of detergent at 50°C. (Detergent 10% Antarox (Igepal) CO-630. Specimen thickness 2 mm. Examination every 4 hours and calculation is based on probability of 50% broken samples.

25 Density - ISO 1183: 1987

[0086]

30

15

20

35

40

45

50

5	4	4	3.	<i>3</i>		2			1:	11	5
		o	5	0		5	9		5	o	
Mechanical data:			934- Ref	934- A	934-B	934- C	934 -D	934 -Y	RM8343	RM8403	ME8152
ESCR	10% F50		33	24	21	25	36	10			
Series 1											
9/1,4 13-10-6											
Tensile properties	Tensile modulus	Мра	620	089	099	650	620	008	585	710	009
	Stress at yield	Мра	18	18	18	18	18	21,5	17,5	20	17
	Strain at yield	%	13	11	12	12	12	11	13	12	13
FWI at-20°C	Max force	N/mm	1430	1390	1400	1410	1410	1470	1425	1490	1310
	Total energy at break	٦	19	19	19	19	19	19	19	20	15
	Total deformation	mm	24	24	24	25	24	23	24	23	22
Density			937,2	937,4					936		933,4
Series 2											
9/1,4 13-30						·					
Tensile properties	Tensile modulus	Мра	630	029	630			755		740	610
	Stress at yield	Мра	18	18	18			21		20,5	17
	Strain at yield	%	12	11	12			11		13	13
FWI at-20°C	Max force	N/mm	1440	1420	1420			1470		1480	1320
	Total energy at break	ſ	19	19	19			19	:	18	16
	Total deformation	mm	24	24	24			23		23	22
Series 3											
4/2 13/30											
Tensile properties	Tensile modulus	Мра	640	002	670	680	645	795	615	705	009
	Stress at yield	Мра	18	18	18	18,5	18	22	17,5	20	16,5
	Strain at yield	%	11,5	11	12	12,5	12	=	12,5	12,5	11
FWI at-20°C	Max force	N/mm	1220	1190	1190	1210	1210	1270	1200	1260	1100
	Total energy at break	ŋ	17	17	17	17	17	17	16	16	13

		ME8152	23	927,5
10		RM8403	24	
15		934- Ref 934- A 934- B 934- C 934- D 934-Y RM8343 RM8403 ME8152	24	935,7
		934 -Y	24	
20		934 -D	24	
5		934- C	24	
	(continued)	934-B	24	6
)	(conti	934- A	25	286
5		934- Ref	24	8'966
,			mm	
5			Total deformation	
o		Mechanical data:		
55		Mechan		Density

11

[0087] 934Y has the most favourable tensile and impact properties but has lower ESCR. Overall, compositions of the invention show improved warpage, stiffness, morphology and comparable or improved ESCR. Grade 934A ha also been found to have increased high temperature stiffness.

### 5 Example 6

[0088] The warpage and thickness data for rotomoulding serial 1 with all blends is displayed in Annex 3 directly and in comparison to RM8343. Data for serial 3 is displayed in Annex 4 directly and in comparison to RM8343. These results are graphically depicted in Figure 2.

[0089] As will be seen, the warpage results for the compositions 934Y, 934A, 934B, 934C and 934 D are better than the results obtained using the available commercial grades.

### Example 7

15

20

25

30

35

40

45

[0090] The following metallocenes are used in this example. Me<sub>2</sub>Si(9Flu)<sub>2</sub>ZrCl<sub>2</sub> (A-metallocene) which produces amorphous attaic polypropylene and rac Me<sub>2</sub>Si(2MeIndenyl)<sub>2</sub> ZrCl<sub>2</sub> (B-metallocene) which produces isotactic polypropylene. These catalysts are available from Boulder.

### **Dualsite catalyst preparation**

[0091] Preparation of catalyst "60/40" in the Table in Example 8 - A pretreated silica carrier, calcined at 600°C was transferred to a small bottle with stirrer bar. The carrier was wetted with toluene (5.5 ml per 2 g carrier). A solution of 2.5 ml of 30% MAO in toluene, 38.4 mg of A-metallocene and 33.4 mg of B-metallocene was prepared and stirred for 30 mins. This solution is added dropwise to 5.5 ml of toluene containing 2 g of silica carrier and stirred for 20 minutes and left overnight. The catallyst is then dried in nitrogen for 2 hours at 40°C.

[0092] The other catalysts in Example 8 were made analogously to "60/40" maintaining the total molar metallocene concentration constant.

### Example 8

[0093] The dry catalyst is fed into a 2L reactor under nitrogen. 650 ml propylene is added to the reactor and prepolymerisation is initiated for 8 minutes at 15°C. The temperature is rapidly raised to 70°C and polymerisation takes place in the absence of hydrogen.

[0094] The reactor conditions are described in the Table below.

Run no	Temp. °C	Time	H <sub>2</sub>	mg cat	weight polymer	Extra TEA, 1M (A54)	Dual site catalyst in mol% (B/A)
6301	70 °C	40 min	0	220	200 g	0,3ml	100/0
6303	70°C	60 min	0	200	15 g	0,1 ml	0/100
6314	70°C	30 min	0	230	190 g	0,25 ml	90/10
6315	70°C	30 min	0	210	190 g	0,25 ml	60/40
6316	70°C	60 min	0	230	130 g	0,25 ml	20/80
6317*	70-80 °C	55 min	0	210	175 g	0,3 ml	20/80

	Mw	Mn	MWD
6301 whole polymer	185 000	82 000	2,3
6301 crystalline phase.	185 000	82 000	2,3
6303 whole polymer	190 000	55 000	3,3
6303 XS phase	185 000	55 000	3,3
6314 whole polymer	190 000	80 000	2,4

50

### (continued)

	Mw	Mn	MWD
6314 crystalline phase.	190 000	80 000	2.4
6315 whole polymer	210 000	80 000	2,6
6315 crystalline phase	210 000	100 000	2,1
6316 whole polymer	240 000	105 000	2.2
6316 crystalline phase	230 000	105 000	2,2

### Results

5

10

15

20

30

35

40

45

50

55

[0095] The properties of the resulting polymers are described in the table below. The polymer components produced form a composition which is expected to be ideal for rotomoulding due to the almost identical Mw and Mn values but differing melting point properties of the isotactic and amorphous components.

[0096] Runs 6316 and 6317 give rise to a polymer composition having xylene soluble fractions of 16 and 19 wt% respectively. (The polymer is boiled in xylene at 137°C for 30 minutes, cooled filtered and the crystalline phase precipitated). The polymer compositions are surprisingly free-flowing powders and it is believed that never before have free flowing polypropylene polymer compositions been prepared having such high xylene soluble fractions. This forms a further aspect of the invention. The high xylene soluble fraction is believed to give rise to a softer polymer which may warp less on rotomoulding.

[0097] Thus viewed from a further aspect the invention provides a free-flowing polypropylene homopolymer powder having a xylene soluble fraction of at least 7 wt%, preferably at least 12 wt%.

[0098] Viewed from another aspect the invention provides a free flowing propylene homo or copolymer powder comprising components A and B wherein:

Component A has a crystalline melting point; and

Component B has a melting point at least 10°C lower than that of component A, preferably 30°C, especially component B is amorphous, has an Mn of at least 25000 g/mol. preferably at least 40000 g/mol and has a comonomer content of less than 20 wt%, preferably less than 5 wt%;

said powder having an Mw of at least 75000 g/mol, a xylene soluble fraction of at least 7 wt%, preferably at least 12 wt%.

[0099] The catalyst used to manufacture said free flowing powder is also new. Thus viewed from another aspect the invention provides a solid, preferably supported multisite catalyst comprising two metallocenes A' and B';

metallocene A' comprising two optionally substituted indenyl groups connection via the 1-position of the indenyl, said metallocene having  $C_s$  symmetry and preferably comprising a group 4A metal;

metallocene B' comprising two optionally substituted fluorenyl groups connection via the 1-position of the fluorenyl, said metallocene B' having  $C_s$  symmetry and preferably comprising a group 4A metal.

Annex

# PREPARATION OF COMPONENTS A & B USING (nBu-Cp), ZrCl,/MAO ON SILICA

Cal temp (°c)/Loading (%)	001/009	600/100	001/009	600/100	600/100	600/100	600/100	001/009
IMP	Dry-mix	Dry-mix	Dry-mix	Dry-mix	Dry-mix	Dry-mix	Dry-mix	Dry-mix
Reac temp	94	94	94	94	85	85	58	58
Reac pres (bar	25,5	25,5	25,5	25,5	23,1	23,1	23,1	23,1
Etene partial pressure (bar)	7,5	7,5	7,5	7,5	7,5	7,5	7,5	7,5
Eten H2 ratio (ppmH2)	420	420	420	420	650	650	650	440
C6/Etene-Cascade (Wt%C8)	0,00	0,00	00,00	00,00	6,00	6,00	6,00	00°9
TOTAL RUN TIME (min)	09	09	09	09	09	60 '	9	09
MFR2 (powder)	6,1	6,4	6,8	6,2	8,1	8,5	8,3	6,5
MFR21 (powder)	65	105	121	103	123,0	122	120	110
FFR (powder)	15,6	16,4	17,8	16,6	15,2	14,4	14,5	16,9
DENSITY (powder kg/dm3	0,9573	0,9577	0,9563	0,9568	0,9224	0,9214	0,9216	0,9245
		COMPO	COMPONENT A			COMP	COMPONENT B	i

EP 1 236 770 A1

ANNEX 2

8-1.4 ald vs 4-2 full cool

Material		1934 A(std)	834_A(std) 834_A(124/2) RM8		343(810) RMB343(104/2) MEB152(810) MEB152(104/2)	EB152(std) h	IE8152(Ic4/2)	Material	Вох	934_A(opt	834_A(opi/934_A(tod   RM8343(o) RM8343(1   ME8152(o   ME8152('ca/2)	3M8343(0)R	M8343(1 M	E8152(o M	E8152('04/	(2)
Warpage								Warpage	$\neg$							
Center		0,45	-0,5		0.48	2	1,92	Center	SZE	0,234375		0,177083		0,541667	=	
		1,61	1,5	2	4.71	1,9	2,5		S3E	0,644		_	1,884	0,76	į	
	SKE	0.67	1.1		1,28	1,86	1,34		S4E	0,5	-0,82838		_	1,238808	-	
	SSE	4.	1.8	.,	4.88	1,71	2,68		SSE	0,528318	0,801504	1,033835	1,827068	0,642857	-	
	SeE	0.31	3.8		5.87	1,8	3,85		SeE	0,075949	0,962025	0,967089	1,488078 (	0,405083	-	
	WAV(\$2-58)E	0,8861	1,058	2,112	3,44	1,682	1,474		WAV(32-	0,358124	0,427648	0,863678	1,380481	0,63945	-	
Max	Szmax	L	0.8		1,02	1,09	1,92	Max	32max	0,338542	0,416667	0,358375	0,53125 (	0,567708	=	
	S3max	1.81	1,5	2.15	4,71	1,9	2,5		S3max	0,644	9'0		1,884	0,78	-	
	S4max	86.0	1,2		1,28	1,79	1,34		S4max	0,514925	0,895522	1,283582	0,955224 1	1,335821	-	
	S5max	7.	1,6		4,80	1,75	2,88		S5max	0,528318	0,801504	_	_	0,857895	-	
	Semax	0.34	3,8		5,87	1,6	3,95		Semax	0,088078	0,982025	_	_	0,405083	-	
	WAv(\$2-S6)m		1,78		3,648	1,626	2,474			0,379143	0,719483	0,916117	1,434115 (	0,657235	-	
Thicknes	1		3,55	3,84	3,62	3,43	3.69	Thicknes		0.98208	0.96208	0.88645	0,98103	0.929539	-	
	28	3.38	3.67		3,73	3,37	3,85		j	0,872727	_	0,932468	0,988831	0,875325	-	
	2C 1	3.49	3.78	3,69	3,89	3,77	4,01		2.C	0,870324	-	0,9202	-	0,94015	=	
L	2.0	3.6	3,83		3,82	3,71	4,05		2.0	0,888889	0,958025	_	_	0,916049	=	
	2 E	3,37		3,58	3,86	3,58	7		2.E	0,8425	0,865	0,895		0,89	-	
	3.4	3,24			3,22	3,28	3,1		3.A	1,045181	0,983871	1,190323	1,03871	1,058085	-	
	3.8	3.23			3,17	3,18	3,18		3.8	1,022152	1,058962	-	1,003185	1,008329	-	
	30	3.52		3,48	3,55	3,68	3,77		3.0	0,933687	0,96617	0,923077	0,841645 (	0,976127	-	
	30	3.55			3.54	3,51	3,67		3.0	0,867302	1,019074		0,964578 (	0.956403	-	
	3.5	3.02			2,58	3,15	2,91		3.6	1,037801	0,886564	1,108529	0,879725	1,082474	-	
	4 A	3.18			3,81	3,38	3,81		4.A	0,83727	0,942257	0,92128	7	0,887139	-	
	48	331	3.78		3,75	3,47	3,67		4.8	0,901907		0,850854	1,021799 (	0,945504	=	
	40	3.57		:   	3,89	3,69	4,08		<b>4.</b> C	0,87831	_		Н	0,908897	-	
	40	3,57			3,81	3,54	3,93		4.0	0,808397	0,951654	-	_	0,900783	=	
	4 E	3.41	3,72	i L	3,88	3,63	3,89		#. m.	0.854637	0,932331		_	0,808774	-	
	5 A	3.14			3.21	3,25	3,53		5.A	0.689518	0,963173	0,929178	0,909348	0,92068	=	
	5.8	32	3.58		3,32	3,32	3,32		5.8	558586'0	1,072289	1,080241	-	-	-	
	50	3.48			3,87	3,67	3,58		5.C	0,972067	0,988827	1,013988	1,02514	1,02514	-	
	50	3.44	3.63	3,53	3,23	3'8	3,37		5.D	1.020772	1,077151	1,047478	j	1,038578	-	
	5.5	3.24			2,74	3,23	2,83		5.E	1,105802	_	1,095563	0,935154	1,102389	-	
	8.4	3.82		3,83	3,73	3,64	3,77		8.A	1,013283		1,015915	_	0,985517	-	
Ĺ	8.8	3.77			4,05	3,54	4,07		8.B	0,92829	-1	0.918918	_	0,869779	-	
	90	3.67		3,63	3,8	3,64	4,03		9 C	0,91067	_	0,800744	0,942929	0.903228	-	
	0.8	3.17	3,61		3,75	3,81	3,75		0.D	1,005333	_		-	1,016	-	
	9 6	3.6				3,42			6.E	1,168831			1,253247	1,11039	_	
	S2 aver	3,474		Ę	6,	3,568			S2 aver	0,889224			_	0,910204	-	
	S3 avar	3,312			3,208	3,38	e.		S3 ever	0,99899		-4	_	1,011439	-	
	S4 aver	3,41	3,702	3,588	3,848.	3,542	C		S4 8ver	0,878158		_	_	0.910072	-	
	SS aver	3.3			3,234	3,384	3,346		S5 aver	0,988252	_ 1	_	-	1,014345	=	
	Se aver	3,726			3,838	3,61	3,74		S8 aver	-				0,985241	-	
	Average	3,4444				3,4948			Average	-1	_	-	_	0,959058	-	
	ThStDev	0,2116435	0,2116435 0,26187338 0.	8 0,17385339	L1	0,18901323		-	ThStDev			0,465982	-	0,608593	=	
	Stdev(av)	0.1730168	0,15458715	9 0,09541488	0,334086012	0,11089274	0,291283367		Stdev(av)	0,693981	0,630711	0,327587	1,148952	0,380704		

ROTO SERINL 1 RESULTS

Meterial		934 Ker 19.	1934 A 19	934 B	37 0	934 D	934 Y	RM8343	RM8403	ME3152	_				7	n 3	5 3	のできるかできた		ر م
Warpage		<del> </del>				Γ					1				}	5			)	•
Center	SZE	1,25	0.43	0.92	0.95	0,42	0	×.0		1.04	Material	Material 8	934 Ref 9	934 A 193	034 8   03	034 C 034 D	D 034 Y	Y (RM8343		RM8403   ME8152
	S3E		5	125	1.5	2.1	22	2.15	2.85	6.	Warbage		-	-	-	-		-	-	L
	SAE	5	190	133	1,35	1.72	142	=	13	1.66	Center	SZE	3 676471	1,323529 7	2,705,882 2	2 794118 1.	1,235294	0	-	O S Chang
	SSE		3	2	2.16	2.47	334	275	196	171		SSE	١.	1	_	<u>.                                    </u>	٠-	1077756	1 1 275481	1
ſ	SBE	1	6	16	222	1,61	0	3.82	24	9	-	SAE	1_	_	_	1	•	C.945667	-	
	WAV(82-5	Γ	989.0	3	1.836	1884	1,382	2312	2,092	1.682		SSE	4-	+-	ŧ₌	٠-	۰	1,214545	133	1.334545 0.62181
Max	SZmax	1	59	15	Ì	990	0.77	690	60	1 09		SGE	0,680628	1_	1		┺	0	1 0 63089	29 O 41AB
	Samax	3.02	191	133	69	2.1	239	2.15	2,85	1.9	-	Warp(S2-	1 028409		0.677083 0	١	┺-	0.6550911	1 0 99053	53 0 74975
	SAMEX	13	0,69	155	15.	1,72	1.42	17.	- 3	1.79	Max	SZmax	1.884058			_		1,115942	1 1304348	_
	SSmax	2.8	=	2	2.16	2,58	Ē	2,3	3,67	1.75		S3max	٠			1	₽	1.111628	11 1375581	At CARTY
	Semax	2,71	0.34	9	22	191	ē.	ē	241	9	-	SAMax		<u>.                                    </u>		_	┺-	0.825581	1 0 889535	
	WAV(32-5		0.838	7	1,708	1,734	1.842	2.284	2277	1.826		SSmax		-	1.	٠.	0.938182 1.2	1214545	1 133/545	
Thickness	×2	l	3.55	345	3.45	350	3.42	364	33.55	343		Samax	0,67581			٠.	-	0,321696	1 0,600998	
	2.8	3,55	338	348	3.36	3.47	3.36	3,53	3,36	337		Warp(32-	0,983218	0,414311	0,660212 0		0,765901 0,8	0,813004	1 1,003534	
Γ	2.5	3,62	3,49	3,63	347	3,56	3.46	3,63	3,62	3,77	Thickness	2.A	0,991758		0.947802 0	0,947832	L	0,93956	1 0.92033	
	2.0	3,53	3,6	3.55	3.77	3,56	3,48	3,63	3,69	3,71		2.8	0,988858	0,835933	0,969359 0	0,935933 0	0.966374 0.9	0.93333	1 0 935933	
	3.5	3,56	3,37	3,53	3,49	3,53	3,38	3,58	300	3,56		2.C	0,98103	<u> </u>	0.98374 0	<u>L</u>	0.984771 0.9	0.937669	1 0 98 (0.3	
	3 A_	3,27	3,24	3,25	3,37	333	3,45	3,69	3.25	3,28		20	0,95664	0,97561	L	1,02168 0		0,943089	22/50	X
	3.8	3,34	3,23	3.45	3,24	337	3,24	3.47	3 13	3.18		2.E	0,994413 0,941341	_	1		_	0.9441341	1 0 988327	27 0 99441
	30	347	3,52	3.43		3.42	3.44	3.48	3,36	3.68		3.A	0,886179 0,876049	_	_			0.924119	1 0 880759	
	3.0	3,54	3,55	3,47		3,54	3,48	3,43	3,56	3.51	L	3.8	0,962536 0,930838	0,930838 (	994236 0	0,933718 0,971182	71182 0.9	0.933718	1 0 902017	
	3 €	3,23	3,02	33		3,22	3.18	3.22	304	3.15		3.0	0,997126	1,011494	0,985632	1,017241 0	0.982759 0.9	0.982506	1 0 982844	
	44	3,35	3,10	3,32		3,32	3,32	3,51	3,48	3,38		3.0	1,03207	1,034985	1,011662	1,058309	1,03207	1,014577	1 1,034985	
	4.8	3,45	3,31	3,46		3,39	3,27	3,49	3,28	3,47		3,6	1,003108		1,021739	1,040373	50	0,967578	1 0,9440	99 0 9782
	Ç.	3,69	3,37	3,71	3,65	3,62	3,48	3,78	3,6	3,69		4.A	0,954416  0,908832	0,908832		1,002849 0	0,945869 0,8	0,945869	1 0,991452	52 0 0826
	4.0	3,68	3,57	3,45	3,54	3,57	3,49	3,56	3,55	3,54		4.B	0,988539 0,948424	0,948424			8,0 715176,0	0,935963	1 0,939628	28 0.6942
	9.E	3,53	3,41	3,45	3,48		3,42	3,61	3,42	3,63		, C	0,978723	0,949468	0,986702 0		0,962766 0,9	0,925532	1 0,930851	51 0,9813
	3.A	3.19	3.14	3,19			3,38	3.28	3,21	3,26		4.0	1,02809	1,002809	0,969101 0		1,007809 0,960337	60337	1 0,897191	91 0,9943
	9.8	3,38	3,2	3,45		3,39	3,25	3,52	3,13	3,32		4.E	0,977839 0,944598	0,944598	955679 0		0,962389 0,947368	47368	1 0,9473	- 8
	ر د د	3,54	3,48	3,56	3,56		3,34	3,63	3,52	3,67		5.A			_		0,9725611 1,030488	30488	1, 0,978655	55 0,8908
	5.0	3,43	3,44	3,39			3,23	3,53	3,45	3,5		5.B				0,931818 0,	0,963058 0,8	0,623295	1 0,8892	35 0,84318
1	3.E	SZ.	3,24	3,25		3,25	3,1	3,21	3.1	3,23		50	_				0,956923 0	0,92311	1 0,969697	
	8.A	3.84	3,02	384			3,77	3,83	3,57	3,64		5.0						132011	1 0,917331	37 0,99150
	6.9	3,86	377	364		3,78	3,66	3,74	3,62	3,54		5.E	_		_			0,965732	1 0,955732	32 1,0067
	6.C	3.73	3,67	374			3,61	3.63	3,7	3,84		8 A	1,002811		1,028721 0		0,994778 0,5	0,984334	1 0,9321	15 0 85039
	,6.D	3,69	3.77	3,64	3,97		3,7	3,72	3,6	3,81		6.8	1,032086			1   \$3686		0,97861	1 0,957914	14 0.94652
	8.E	3,42	3.6	3,38			3,39	3.2	3,28	3,42		90	1,027548			1,002755 0,991738	30 82718	0,966342	1,019284	
	S2 aver	3,574	3,474	3,528			347	3,638	3,492	3,568		90	0.991935			1,087264 0,939247		0,994524	1 0,957742	42 1,02419
	S3 aver	3,37	3,312	3,378		3,376	3,35	3,458	3,264	3,36		8.E	1,05875		1,05	1,084375 1,065625		1,059375	1 1,01875	
	S4 aver	3,534	3,41	3.478			3,356	3 558	3.446	3.542		52 8781	0,982403		0,959764 0	0,952764 0,964268 0,973612		0.940077	1 0,959868	57082,0 33
	SS aver	3,366	?	3358	╝	1	3,272	3434	3,282	3,394		S3 BVer	0,974552		976865	0,976865 0,990748 0,976287	76287 0.5	0,968768	1 0,9438	38 0,9716
	S6 ever	3,708	3,726	3.664	3,696	-	3,606	3,624	355	1,61		S4 BVer	0.082499		0,869283	O REGRESS 0,983268 0,969325 0,947018	969325 0.5	47018	1 0,960959	50 0,987
	-	3,5104	3,444	3,4632	- 1	3,4756	_		3,4088	3,4946		SS ave	2.98019B	_1	0,98078	_	969132 0.9	0.952825	1 0,955737	37 0,98835
		0,179153 0,211544 0,170898	0.211544	0,170898	- 1	0,160833	0,153577			0,189013		Se aver	1,023179		_4	1,019268	1,00883 0,9	0,995033	1 0,979581	11 0,59613
	Stdev(av)	0 145056	0.1730:7	0,121512	0,104638	0,131015	0,123843 0,095415	0.095415	0,127739	0,110893		Average	0.989432	_		0,892897 0,979594	79594 0,5	0,960767	1 0,960203	23 0,93500
	•						-	-		-	-	Thistoer	1.030481	1.215793	0.9929991	0.95725 0.025109	225100 0 0	O RELITA	1 1 084736	36 1 08719
															1			2		

									5.5	-	E																						.3		200		
	÷	NEB152	1	0.530795 1.048475	\$2.53 \$2.53 \$3.53	0 7 10 11	1 682353	0.510788	25/2/20	0,077013	152630	NAME OF THE PERSON OF THE PERS	1012177	00000	1003163	1,010/07	0 000	1091972	52,550	200	0,978967	043702	1078351	1.009588	0.075477	20,240	1,089341	1,010724		e Constitution	(tolst)	1010101	38	8	0.074486	1000	67,875
5		RM8403 N	1797867	22.615	1202	1 185247 0 672	1,7759		127657	1,185247	9717	1077-44 10194	005367	90500	0043878	2	रङ	1011266	200	200	78.0	2200	0,912371			-	0.934307	1827.23	11188	200	0,589119 0,79792	0.989485	86.836	238336	3563780	0.985.518 1.0160	1,010756 0,001 1876
	343	RN.8343 R	F			#	F		1	٦	- 10/12	T NAV	-		3		=	3	T	1		T		=		1			7	-	7	7	1	-	ŝ	=	#
10	RM 6343	П	58333	\$378175	B\$\$\$\$\$0	0450966	70528	200554	95559	59865	_	1001100	59785	13162	70	769	1,146265	1182001	0806107	44827	3 0949333 0,000333	20102	551.6	888	05458	1,114551	1,10218	277561	27077	71 10065138 1.007687 1.018687	110533	2175	13217	0185531	0,914018	0,00000	63759
	5	D Bld V		1,367,188 5.3	\$ 0 58850	0 668073 0	1 2176	0,300034 0,509554	2865 2885 985 985	77385 0.4	0.501681 0.723788 0.808908 0.669509 0.764574	07238104	7681 05	4200 0	5 0 964 266 10 977041	1578 0	0,995691		0 10 10	98130	19333 0	0 20 00	0.956186 0.991753	1,024077 1043814 1,05919	200	1.000788	87044	21743 0	3	1,840	36781 0	7387	100500	2008	4765	0 986516 0	0.443542 0.588656 0.18.862 0.838548 0.588372 0.65355 0.731543 0.462714 0.470871 0.580213 0.580815 0.453759
15	CORPACES	0 PRO	45 200	40025 1.30			117647 16	0100	4879 045	8998 0.80	300	0.991713 0.997238	1,02561 1,00	003144 09	0.201245 0.00	1037767 0 0	9937 098		01	2125 0.0	0 865333 00	0001111	3709 0.0	01/2/20	275 09	6576 1.0	10,881752 1,082044	23.10 0.8	600		30 9539	5604 0.9	1024605	6070	0,208286 0,917144 0,944767	0.083605 0.0	27 05
15		9 <u>7</u>	115 4.5	575 1.4	\$ 0 13	278 0 835308	1~	001057 O 450100	247 055	529 0 83	000 917	100		20 00	155 099	366 1037267	937 1,059937	352 09	1070000 1.110160	27	380 [11		0658)63 0 353402 0 943769	3	027 0 50	30031 101	004821 0,88	718 0.0		80	139 0.76	856 0.89		101 853	280 081	0000	871 0.58
	RESULTS	87.8	3315	13 0.921875	18 0.545.567	57 O 736478	1,500,335	1000	16 0 380247	50 0.870	10.72	0 840653 1 1006787	0 853914 1,005362	1660 18	1012755	0 95472 0 947705 0 981388	28 1059			57 0 28 16 27		21 0 02697	53 0 53	000	76 0 670027	39 1,130	7 1004	72 0 884718	000	0.952857	34 0765	57 0 536858	1002230	1,08	100 0 50		14 9.470871
20	RE	Y PE	1301507	6 0.867183	0.379718	0.494767		의	6 0 329218	1 0.0473	0.050	90930	5 0.888	38 0.965	261000	2000	1 1053078	2 1 0781	1 4444 1 1056497	8 09477	10.03-335 1.002887		2 0 55 87	1,05019	0.893733 0.064578	4 1.738	1,098	12 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2	772880	0 0 0 0 0	1 0 6 73 834	의		1,0\$8998	3 0.918700		12 0 de 2774
		3	50100	297658	0.05.727			9000	0.691588	0,8160141		1.02/524	1013405	103.13	08.40		1.07571	100:072	2705270	0.95275	0.03.33	N X X X	0.930412	1,10289	0.19373	0.950464	1,1058	0014200	178830	7	Н	7	1000	0.099145	0.930693	0.007268	SELLO SELLO
25		Ful cookin	STE	77	SSE	WerdS2	Sames	Sales Sales	Sympa	Series	Yents	Z	7.9	37	270	_	8	ပ္	0	4	87		4	<b>1</b>	200 200 200 200 200 200 200 200 200 200	35	SE	3	8	9	35	Szerr		55 gree	38 Per	ACCUPANT OF	STOWER
	4	Mesand	. 1	T			Mer	1			T	1							1																		
20	ANNEK	H	131	200	180	1	Ţ	157	1		120	1	<u>'1'</u>	101	405	1	100		367		301	9	188	355		333	282	111	107	1	33	193	3.72		11	3544	0.25125
3 <i>0</i>	<b>a</b>	3 MEB152	11		11	ľ	L	1	575	1			3,5			1	1	П		1	150	1	ı	П			L	123	369	] 3;	3	3,764		1	Ш		
		RN48403		9	11	1	Ĺ	i ł		1	1 1		1			317	L	П		1		1		П	1	1	1	11				11		ł	3,818	17	0
35		RMB345		474	Ц		L	П	L	L	Ц	1	1	П	J		L	П		1		1	L	Ш	1		L			$\perp$	L	Ц		1		3 5884	0.334039
		7 78	2.	236	П	-27			1	L	2712	1	158	Ц		1	175	П		8 5	1	1				1		Ц	Ц		L	Ц		725		3433	0.151586
40	SINGAU	O Mg	600	20 52	241	1	105	17	140	15	2374	**	374	3.78	3.78	3.67		347	3.05	7	3.56	88		335		378	ā	355	187		150	3.14	CC .	100	3.676	35384	0,162,149
		934 C	218	7	2	5	2.16	2,12		15	207	090	12	3.4	187			3.42	3,93		362	7	308	378	35	23	2.63	3,63	103	9	28.	3.75	3386	346	355	3,5276	0.193842
45	SERIAL	8 B 78	1.62	===	265	115	162	117	1,18	1	2.562	1	375	988	3.67		3.36	ž	724		3,92		235	35	3 5	365	6	33	350	1	3	3,754	3382		3,488	3,5892	0,157312
45	ROTO SERIAL 3	184 A 181	55	=	2	7	80	51		300	13	+	367	326	388	200	12	100	3	105	3.76	1	377	76	22	195	301	335	38		3.18	3,644	3336		3,576	3,3312	0,154597 0
•	en en	034 Ref 1934	177	*		4.70	301	32		4.79	3642	133	378	323	382	1		377	3	150	368		186	250	27.	300	333	341	292	3 2	160	391	1	1278	157	3,5228	322045 0. 0.2444 0.
50		P	H	+		763	ļ	¥	E E	¥	Wertst	$\dagger$		H	+	+	-	H	1	+	H	$\dagger$	+	H	$\dagger$	+	-		+	Ţ	Π	Ę	Т	Т	28.5%	Aver 8go	STOW 0.
			4 1	3,3	SSE.	1	Es	ES.	E SC	Sem	We	Thekines 2A		32	2	1	3.8	3.0	9:	1	48	1	1	¥\$		25	SE	¥ 9	83	1	9E	32.	B	38	155	AW	調
55		Material	) (4)			1	Ę		1			Ë		Ц	1				Ţ			Ĺ,	Ц	1	L						Ĺ		1	1	Ц		Ш

### Claims

5

10

15

20

25

30

1. A polymer composition suitable for rotomoulding comprising

I) an ethylene homopolymer or copolymer with at least one other  $C_{3-10}$   $\alpha$ -olefin, having a melt flow rate of 0.5 to 30, a molecular weight distribution (Mw/Mn) of less than 4, an Mw of 50,000 to 110,000, a density of 0.940 g/cm<sup>3</sup> to 0.970 g/cm<sup>3</sup> and a melting point of 100 to 145°C;

OR

I) a propylene homopolymer or copolymer with at least one other  $C_{2-10}$   $\alpha$ -olefin, having a melt flow rate of 0.5 to 30, a molecular weight distribution (Mw/Mn) of less than 4, an Mw of 150,000 to 300,000, and a melting point of 100 to 170°C;

and

II) an ethylene homo or copolymer with at least one other  $C_{3-10}$   $\alpha$ -olefin, having a melt flow rate of within 40% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4, an Mw of within 30% of the Mw of component (I), a density of 0.880 g/cm³ to 0.940 g/cm³ said density being at least 0.010 g/cm³ less than the density of component (I) and a melting point of at least 5°C less than that of component (I);

OR

II) a propylene homo or copolymer with at least one other  $C_{2-10} \alpha$ -olefin having a melt flow rate of within 40% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4, an Mw of within 30% of the Mw of component (I), and a melting point of at least 10°C less than that of component (I).

2. A polymer composition as claimed in claim 1 comprising

I) an ethylene homopolymer having a melt flow rate of 0.5 to 30, a molecular weight distribution (Mw/Mn) of less than 4, an Mw of 50,000 to 110,000, a density of 0.940 g/cm<sup>3</sup> to 0.970 g/cm<sup>3</sup> and a melting point of 100 to 145°C:

and

II) an ethylene copolymer with at least one  $C_{3-10}$   $\alpha$ -olefin having a melt flow rate of within 40% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4, an Mw of within 30% of the Mw of component (I), a density of 0.880 g/cm<sup>3</sup> to 0.940 g/cm<sup>3</sup> said density being at least 0.010 g/cm<sup>3</sup> less than the density of component (I) and a melting point of at least 10°C less than that of component (I).

- A composition as claimed in claim 2 wherein component (II) is an ethylene copolymer with butene, hexene or octene.
  - 4. A composition as claimed in claim 3 wherein component (II) is an ethylene copolymer with hexene.
- 40 5. A composition as claimed in any one of claims 2 to 4 wherein the melt flow rates of components (I) and (II) are 4 to 10.
  - 6. A composition as claimed claims 5 wherein the melt flow rates of components (I) and (II) 6 to 8.
- 7. A composition as claimed in any one of claims 2 to 6 wherein the molecular weight distribution of the composition is less than 4.
  - A composition as claimed in any one of claims 2 to 7 wherein the density of component (II) is 910 g/cm<sup>3</sup> to 0.930 g/cm<sup>3</sup>.
- A composition as claimed in any one of claims 2 to 8 wherein the density of composition is in the range 0.925 to 0.950 g/cm<sup>3</sup>.
  - 10. A composition as claimed in claim 9 wherein the density of composition is in the range 0.930 to 0.940 g/cm<sup>3</sup>.
- 11. A composition as claimed in any one of claims 2 to 10 wherein the melting point of component (I) is in the range 125°C to 135°C and the melting point of component (II) is in the range 100°C to 125°C.
  - 12. A composition as claimed in any one of claims 1 to 11 wherein the molecular weight distribution each of components

(I) and (II) is less than 3.

5

10

20

25

30

35

40

45

50

55

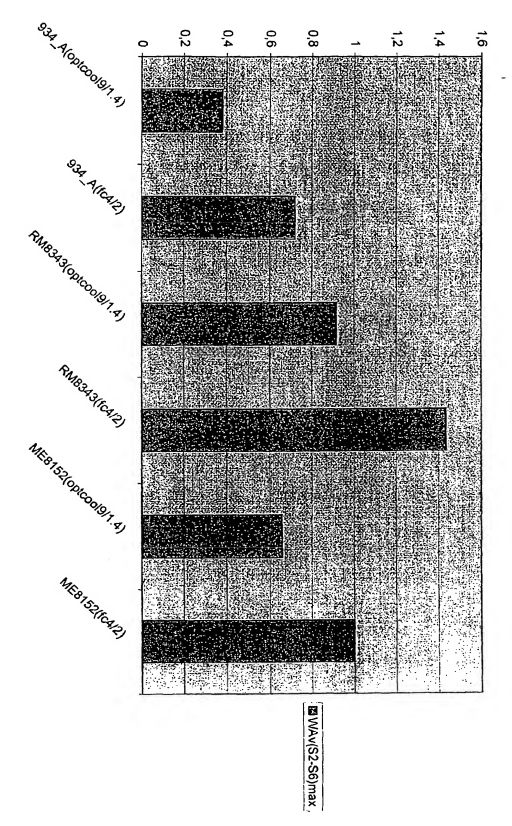
### 13. A composition as claimed in claim 1 comprising

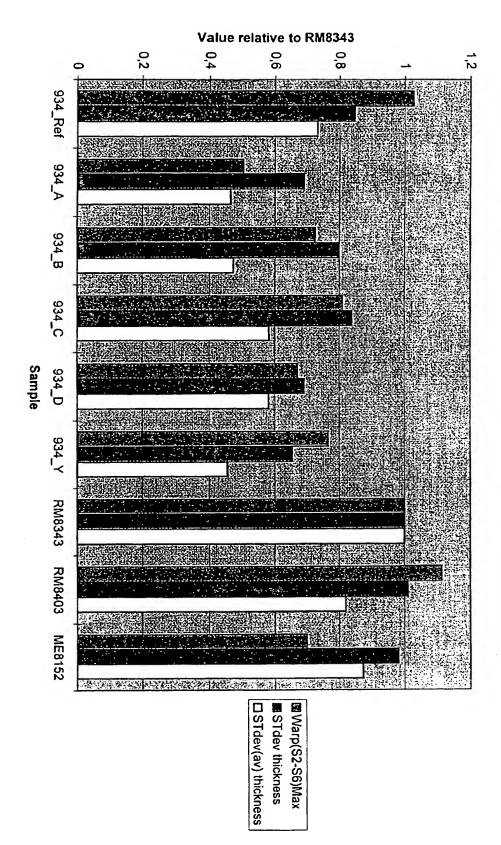
I) a propylene homopolymer having a melt flow rate of 4 to 10, a molecular weight distribution (Mw/Mn) of less than 4, an Mw of 150,000 to 300,000, and a melting point of 100 to 170°C; and

II) a propylene homopolymer having a melt flow rate of within 40% of the melt flow rate of component (I), a molecular weight distribution of (Mw/Mn) of less than 4, an Mw of within 30% of the Mw of component (I), and a melting point of at least 10°C less than that of component (I);

wherein said composition has a molecular weight distribution (Mw/Mn) of less than 4.

- 15 14. A composition as claimed in claim 13 wherein component (II) is amorphous.
  - 15. A composition as claimed in any one of claims 1 to 14 wherein the ratio of component (I) to (II) is from 4:1 to 1:4.
  - 16. The use of a polymer composition as claimed in any one of claims 1 to 15 in rotomoulding.
  - 17. A process for the preparation of an article comprising rotomoulding a composition as claimed in any one of claims 1 to 15.
  - 18. A process as claimed in claim 17 wherein retemoulding is effected at a retation speed of 9/1.4 RPM; heating for 13 minutes in oven at 270°C; fan assisted cooling for 10 minutes followed by 6 minutes ambient cooling in the absence of a fan.
    - 19. An article comprising a polymer composition as claimed in any one of claims 1 to 15.







### **EUROPEAN SEARCH REPORT**

Application Number

EP 01 30 1873

ategory	Citation of document with indica	tion, where appropriate.	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
X	US 6 111 023 A (CHUM F AL) 29 August 2000 (26 * column 9, line 4; ex	100-08-29)	1-12, 15-19	C08L23/10 C08L23/16
X	DATABASE WPI Derwent Publications L AN 1996-045404 XP002172829 & JP 07 309909 A (MITS 28 November 1995 (1995 * abstract *	td., London, GB;	1-12, 15-19	
A	AU 21280 99 A (J.R.COL 30 September 1999 (199 * example 1 *		1-12, 15-19	
				TECHNICAL FIELDS SEARCHED (Int.C1.7)
				C08L
	•			
	The present search report has been	drawn up for all claims		i 
	Place of search	Date of completion of the search		Examine:
	THE HAGUE	7 August 2001	SCH	MIDT, H
X : part Y : part docu A : tech	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone icularly relevant if combined with another iment of the same category inological background -written disclosure rimediate document	E : eadier patent after the filing D : document cite L : document cite	siple underlying the document, but publicate so in the applications of for other reasons as same patent family	shec on, or



Application Number

EP 01 30 1873

CLAIMS INCURRING FEES
The present European patent application comprised at the time of filing more than ten claims.
Only part of the claims have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims and for those claims for which claims fees have been paid, namely claim(s):
No claims fees have been paid with n the prescribed time limit. The present European search report has been drawn up for the first ten claims.
LACK OF UNITY OF INVENTION
The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:
see sheet B
All further search fees have been paid within the fixed time limit. The present European search report has been drawn up for all claims.
As all searchable claims could be searched without effort justifying an additional fee, the Search Division did not invite payment of any additional fee.
Only part of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the inventions in respect of which search fees have been paid, namely claims:
None of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the invention first mentioned in the claims, namely claims:
1, 5, 6, 7, 11, 12, 15 - 19 partially, 2, 3, 4, and 8 - 10



## LACK OF UNITY OF INVENTION SHEET B

**Application Number** 

EP 01 30 1873

The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:

1. Claims: 1, 5, 6, 7, 11 ,12, 15-19 partially, 2, 3, 4, and 8-10

as far as compositions of polyethylene I) with polymer II) are concerned

2. Claims: 1, 5, 6, 7, 11, 12, 15-19 partially 13, 14

as far as compositions of polypropylene I) with polymer II) are concerned

### ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 01 30 1873

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

07-08-2001

9-08-2000	US US US US US AT AU	5677383 A 5272236 A 5278272 A 5847053 A 6316549 B	14-10-1997 21-12-1993 11-01-1994 08-12-1998 13-11-2001
	US US US US AT	5272236 A 5278272 A 5847053 A 6316549 B	21-12-1993 11-01-1994 08-12-1998
	US US US AT	5278272 A 5847053 A 6316549 B	11-01-1994 08-12 <b>-</b> 1998
	US US AT	5847053 A 6316549 B	08-12-1998
	US AT	6316549 B	
	AT		
		174945 T	15-01-1999
		676047 B	27-02-1997
	AU	6638894 A	21-11-1994
	BR	9406668 A	23-01-1996
	CN	1121730 A	01-05-1996
	CZ	9502830 A	15-05-1996
	DE	69415518 D	04-02-1999
			20-05-1999
			23-08-1999
	EP		14-02-1996
	ES		01-03-1999
	FI	955148 A	27-10-1995
	HK	1014724 A	14-07-2000
	HU	72909 A,B	28-06-1996
	JP	2894517 B	24-05-1999
	JР	8509767 T	15-10-1996
	NO	954327 A	22-12-1995
	NZ	265562 A	24-10-1997
	PL	311306 A	05-02-1996
	RO	116409 B	30-01-2001
	SI	9420032 A	30-04-1996
	WO	9425523 A	10-11-1994
			26-10-1995
			29-04-1993
	• • • • • • • • • • • • • • • • • • • •		25-09-1997
			25-03-1999
			03-07-1997
			20-11-1997
			04-03-1999
			02-06-1999
			03-08-1994
	_		09-07-1997
			03-03-1999
			03-03-1999
			01-10-1997
			01-04-1999
			31-05-1994
	-		12-10-1999
	-		19-01-1995
	KR		15-07-2000
	KR	263803 B	16-08-2000
	al Journal of the Euro	DE DK EP ES FI HU JP NNZ PRO NZ CA DE DE DE EP ES FI JP KR KR	DE 69415518 T DK 696300 T EP 0696300 A ES 2125452 T FI 955148 A HK 1014724 A HU 72909 A,B JP 2894517 B JP 8509767 T NO 954327 A NZ 265562 A PL 311306 A RO 116409 B SI 9420032 A WO 9425523 A ZA 9402902 A CA 2120766 A DE 9219090 U DE 9219173 U DE 69220077 T DE 69228265 D DE 69228265 T EP 0608369 A EP 0783006 A EP 0899278 A EP 0899278 A EP 0899279 A ES 2103976 T ES 2127030 T FI 941727 A JP 2963199 B JP 7500622 T KR 262024 B

### ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 01 30 1873

This annex lists the patent family members relating to the paten; documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way hable for these particulars which are merely given for the purpose of information.

07-08-2001

	Patent document ed in search rep:		Publication cate	Patent family member(s)	Publication date
US	6111023	A		US 5380810 A US 5427807 A US 5562958 A US 5395471 A	27-06-1999 08-10-1996
JР	7309909	Α	28-11-1995	NONE	
AU	2128099	Α	30-09-1999	NONE	

For more details about this annex : see Official Journal of the European Patent Office, No. 12/82

V